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# The Role of Exo1p Exonuclease in DNA End Resection to Generate Gene Conversion Tracts in *Saccharomyces cerevisiae*

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**ABSTRACT** The yeast Exo1p nuclease functions in multiple cellular roles: resection of DNA ends generated during recombination, telomere stability, DNA mismatch repair, and expansion of gaps formed during the repair of UV-induced DNA damage. In this study, we performed high-resolution mapping of spontaneous and UV-induced recombination events between homologs in *exo1* strains, comparing the results with spontaneous and UV-induced recombination events in wild-type strains. One important comparison was the lengths of gene conversion tracts. Gene conversion events are usually interpreted as reflecting heteroduplex formation between interacting DNA molecules, followed by repair of mismatches within the heteroduplex. In most models of recombination, the length of the gene conversion tract is a function of the length of single-stranded DNA generated by end resection. Since the Exo1p has an important role in end resection, a reduction in the lengths of gene conversion tracts in *exo1* strains was expected. In accordance with this expectation, gene conversion tract lengths associated with spontaneous crossovers in *exo1* strains were reduced about twofold relative to wild type. For UV-induced events, conversion tract lengths associated with crossovers were also shorter for the *exo1* strain than for the wild-type strain (3.2 and 7.6 kb, respectively). Unexpectedly, however, the lengths of conversion tracts that were *unassociated* with crossovers were longer in the *exo1* strain than in the wild-type strain (6.2 and 4.8 kb, respectively). Alternative models of recombination in which the lengths of conversion tracts are determined by break-induced replication or oversynthesis during strand invasion are proposed to account for these observations.

THE 5'–3' exonuclease Exo1p was first described in *Schizosaccharomyces pombe* (Szankasi and Smith 1992) and has been extensively characterized in *Saccharomyces cerevisiae* (Tran *et al.* 2004). Exo1p is involved in multiple cellular pathways, including DNA mismatch repair, telomere processing in response to telomere damage, and DNA lesion bypass. Its most relevant function to the research described below is in the processing of broken DNA ends. Following double-strand break (DSB) formation in meiosis and mitosis, the broken ends are processed by degradation of the 5' ends. For ends generated in mitosis, this resection process

is initiated by removal of a small number of nucleotides by Sae2p and Mre11p/Rad50p/Xrs2p, followed by more extensive resection by two pathways, one involving Exo1p and the other utilizing Sgs1p/Top3p/Rmi1p and Dna2p (Mimitou and Symington 2008; Zhu *et al.* 2008). Strains lacking Exo1p are still capable of extensive resection of DNA ends, although the rate of resection is slower than in wild-type strains.

The effect of Exo1p on the frequency of meiotic and mitotic recombination has been examined in several studies employing different methods with variable results. Diploids that lack Exo1p have a reduced frequency of meiotic crossovers and reduced gene conversion at some loci (Kirkpatrick *et al.* 2000; Tsubouchi and Ogawa 2000). The extent and/or rate of end processing are reduced in both meiotic (Tsubouchi and Ogawa 2000; Keelagher *et al.* 2011) and mitotic (Mimitou and Symington 2008; Zhu *et al.* 2008) cells. In mitotic cells, loss of Exo1p lowered the frequency of single-strand annealing up to sixfold in some genetic assays (Fiorentini *et al.* 1997), but had little effect in physical assays of

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single-strand annealing (Mimitou and Symington 2008; Zhu *et al.* 2008). Loss of *Exo1p* elevated the frequency of gap repair as detected in a plasmid assay (Symington *et al.* 2000; Guo and Jinks-Robertson 2013). Part of this elevation reflects *Exo1p*-mediated destruction of the transforming plasmid (Guo and Jinks-Robertson 2013). In summary, in most assays of recombination, loss of *Exo1p* has a relatively small effect, presumably because of the functional redundancy in end processing represented by the *Sgs1p/Top3p/Rmi1p* and *Dna2p* pathway. Most previous studies of the effects of the *exo1* mutation on mitotic recombination were done using duplicated genes or plasmid–chromosome recombination in haploids. In our analysis, we examine the effects of *Exo1p* on spontaneous and UV-induced events occurring between homologs in a diploid.

In addition to its role in end resection, *Exo1p* is involved in expanding the 30-nucleotide gaps that remain following the removal of UV-induced pyrimidine dimers (Giannattasio *et al.* 2010). Giannattasio *et al.* proposed that *Exo1p* (in competition with DNA polymerases that fill in the 30-base gaps) can expand the short gaps to ~1 kb, triggering the G<sub>1</sub>/S checkpoint. Below, we show that *exo1* strains have reduced levels of UV-induced recombination events.

The frequency and map positions of recombination events were examined in a diploid heterozygous for ~55,000 single-nucleotide polymorphisms (SNPs) distributed throughout the genome (Lee *et al.* 2009). Crossovers and gene conversion events occurring between homologs can be detected as loss of heterozygosity (LOH), using oligonucleotide-containing microarrays (described further below). Some of the patterns of LOH observed in previous studies (St. Charles *et al.* 2012; St. Charles and Petes 2013; Yin and Petes 2013) are shown in Figure 1.

If a single chromatid is broken in S or G<sub>2</sub>, the resulting crossover leads to LOH from the position of the DSB to the end of the chromosome; the two daughter cells (D1 and D2) have reciprocal patterns of LOH (Figure 1A). Crossovers in mitosis are frequently associated with gene conversion, a region of DNA transferred nonreciprocally between the two homologs (Lee *et al.* 2009). The conversion tract (outlined in blue in Figure 1) results in the two daughter cells having a switch from heterozygous SNPs to homozygous SNPs at slightly different transition points with the difference defining the length of the conversion tract. Within the boxed region in Figure 1A, since three of the four chromosomes in the daughter cells have the “red” form of the SNP and one has the “black” form, this type of conversion is called “3:1.” Below, we describe 3:1 conversion events as single-chromatid breaks (SCBs), as presented previously (Yin and Petes 2013). As shown in the right side of Figure 1, we can depict the two chromosomes in each daughter cell as a single line with green, red, and black segments indicating heterozygous SNPs, SNPs homozygous for the red allele, and SNPs homozygous for the black allele, respectively.

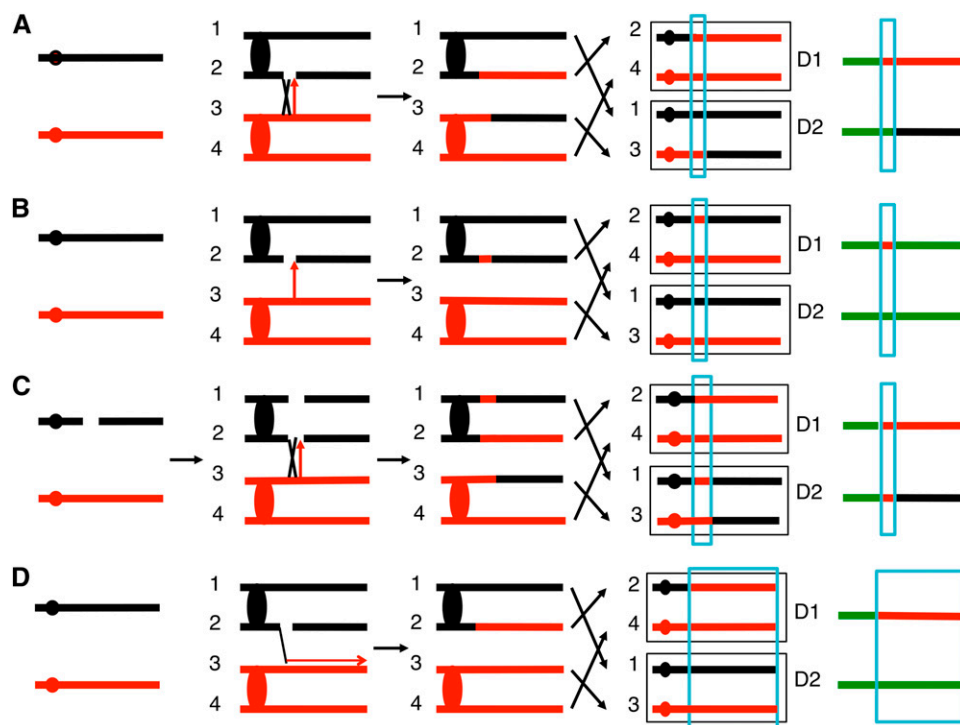
Figure 1B shows the pattern of interstitial LOH generated by a 3:1 gene conversion unassociated with a crossover. Pre-

viously, we found that spontaneous crossovers were frequently associated with 4:0 conversion tracts (Lee *et al.* 2009; St. Charles *et al.* 2012; St. Charles and Petes 2013). In this class of events (Figure 1C), both daughter cells have a chromosome region that is homozygous for the same allele (the red allele in Figure 1). This pattern requires the repair of two sister chromatids broken at the same position, and we suggested that chromosome breakage in G<sub>1</sub> of the cell cycle, followed by DNA replication, would produce such a situation (Lee *et al.* 2009). We also observed conversion tracts that contained a 3:1 section adjacent to a 4:0 section (3:1/4:0 hybrid tracts). These hybrid tracts are also likely to represent the repair of two broken sister chromatids with the extent of the repair being different for the two events. Both 4:0 and hybrid tracts are called double-sister chromatid breaks (DSCBs). Among spontaneous crossovers, about two-thirds were DSCBs, and about half of UV-treated cells (15 J/m<sup>2</sup> treatment of synchronized G<sub>1</sub> cells) had DSCBs. Thus, in *S. cerevisiae*, a substantial fraction of events that lead to LOH reflect DSCBs rather than breaks made during DNA synthesis (Lee and Petes 2010).

Another pattern of LOH is that produced by break-induced replication (BIR) (Figure 1D). In this mechanism, the chromosome fragment located centromere distal to the break is lost. The end of the centromere-proximal fragment invades the homolog, copying its sequences to the chromosome terminus (Llorente *et al.* 2008). As a consequence of BIR, one daughter cell has an LOH event extending to the telomere, and the other remains heterozygous.

To distinguish among the patterns of LOH shown in Figure 1 and to map the positions of the LOH, we used diploids generated by crossing two sequence-diverged haploids, W303a and YJM789 (St. Charles and Petes 2013; Yin and Petes 2013). This diploid, in addition to being heterozygous for SNPs located throughout the genome, contained markers on either chromosome V (YYy29.8) or chromosome IV (YYy34) that allowed identification of reciprocal crossovers (Figure 2). The diploids are homozygous for *ade2-1*, an ochre mutation. In the presence of zero, one, and two copies of the ochre suppressor *SUP4-o*, the diploids form red, pink, and white colonies, respectively (Barbera and Petes 2006). The *SUP4-o* gene is integrated near the telomeres of chromosomes V (YYy29.8) and IV (YYy34) in the strains used in our study. In these diploids, crossovers between the centromere and the heterozygous *SUP-o* marker produce red/white sector colonies. Only half of the crossover events are detectable by this system (Chua and Jinks-Robertson 1991), since cosegregation of two recombined chromosomes into one daughter cell and two unrecombined chromosomes into the other daughter does not lead to LOH.

Crossovers identified as red/white sector colonies were subsequently mapped by SNP microarrays (St. Charles *et al.* 2012). Each of 13,000 SNPs is represented by four 25-base oligonucleotides, two identical to the Watson and Crick strands of the W303a-derived SNP and two identical to the Watson and Crick strands of the YJM789-derived SNP.



**Figure 1** Patterns of LOH resulting from crossovers, BIR events, and gene conversions. The homologs derived from W303a and YJM789 are shown in red and black, respectively. Ovals and circles indicate centromeres. In A–D, the initiating DSB is on the YJM789-derived homolog. The green, red, and black lines on the right summarize patterns of heterozygosity and homozygosity in the daughter cells after the recombination event. In this depiction, green, red, and black signify heterozygosity, homozygosity for SNPs derived from the W303a homolog, and homozygosity for SNPs derived from the YJM789 homolog, respectively. (A) Crossover initiated with an SCB associated with a 3:1 gene conversion. The blue rectangle outlines the conversion event. (B) Gene conversion (3:1) unassociated with a crossover, initiated by an SCB. (C) Mitotic crossover and an associated 4:0 conversion event. Replication of a chromosome broken in  $G_1$  results in two broken sister chromatids. The repair of this DSCB (one repair event associated with a crossover) generates the observed LOH pattern. (D) BIR event initiated by an SCB.

If a SNP is heterozygous in genomic DNA isolated from a red or white sector, it will hybridize to the SNP-specific oligonucleotide to about the same extent as a control heterozygous diploid. LOH is detected by an increased ability of the genomic DNA isolated from a sector to hybridize to the SNPs identical to one parental haploid (W303a or YJM789) and a decreased ability to hybridize to the SNPs of the other parental strain (other details in *Materials and Methods*). This analysis was used to examine spontaneous and UV-induced recombination events in the *exo1* diploid, allowing us to define the location of recombination events, the types of gene conversion tracts, and the length distribution of conversion events. We show that the loss of *Exo1p* reduces the rate of UV-induced, but not spontaneous, recombination events and that the lengths of crossover-associated conversion events are reduced in the *exo1* strain. We also show that a hotspot for spontaneous mitotic recombination events associated with very long conversion events in wild-type cells is absent in the *exo1* diploid.

## Materials and Methods

### Strains

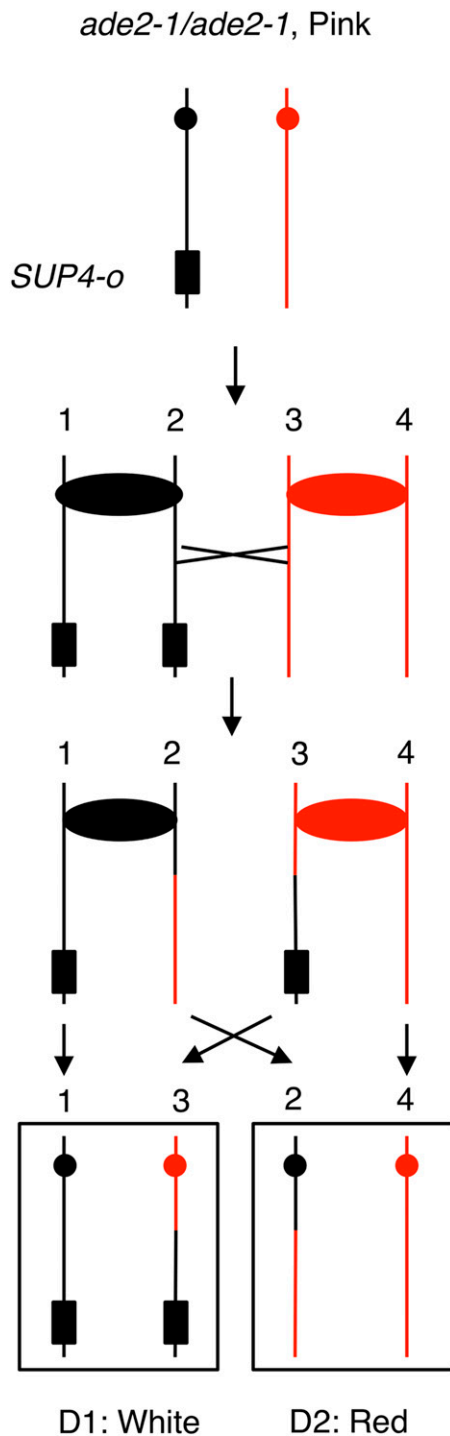
Data comparisons from four isogenic diploid strains were used in our study. All strains were derived from crosses of the two sequenced-diverged haploids W303a and YJM789 (Lee *et al.* 2009) and were deleted for the *MAT $\alpha$*  to allow their  $G_1$  synchronization with  $\alpha$ -factor. The diploids PG311 and YYy29.8 had a heterozygous *SUP4-o* marker near the left telomere of chromosome V, and JSC25 and YYy34 had *SUP4-o* located near the right telomere of chromosome IV.

As explained in the text, the *SUP4-o* marker allows the detection of crossovers.

The wild-type diploid PG311 has the genotype *MAT $\alpha$ ::NAT ade2-1/ade2-1, can1-100/can1::SUP4-o GAL2/gal2 his3-11,15/HIS3 leu2-3,112/LEU2 RAD5/RAD5 trp1-1/TRP1 ura3-1/ura3 V9229::HYG/V9229 V261553::LEU2/V261553* (Lee *et al.* 2009). The *exo1* diploid YYy29.8 is isogenic with PG311 except that the *MAT $\alpha$*  gene in the YYy29.8 strain was disrupted with *KANMX* instead of *NAT*. The wild-type diploid JSC25 has the genotype *MAT $\alpha$ /MAT $\alpha$ ::HYG ade2-1/ade2-1 can1-100::NAT/CAN1::NAT GAL2/gal2 his3-11,15/HIS3 leu2-3,112/LEU2 RAD5/RAD5 trp1-1/TRP1 ura3-1/ura3 IV1510386::KANMX-can1-100/IV1510386::SUP4-o* (St. Charles and Petes 2013). YYy34.2 and YYy34.9 are isogenic *exo1* derivatives of JSC25 except that the *MAT $\alpha$*  gene in the YYy34 strains was disrupted with *URA3* instead of *HYG*. The details of the constructions of YYy29.8 and YYy34 are in *Supporting Information, Table S6*.

### Media and genetic methods

Standard YPD media were used for all experiments except for the sectoring assays that utilized MAB6 solid medium (SD-Arg with 10 mg/ml adenine). For the UV-induced sectoring experiments, cells were synchronized in  $G_1$  by  $\alpha$ -factor as previously described (Lee and Petes 2010). After removing the  $\alpha$ -factor, cells were immediately plated on MAB6 plates and irradiated with 15 J/m<sup>2</sup> of ~254 nm UV, using a TL-2000 UV Translinker. The plates were covered with foil to prevent photoreactivation and incubated for 2 days at 30°. To examine the frequency of spontaneous sectors, we suspended single colonies grown on YPD plates in



**Figure 2** Colony-sectoring assay for detecting mitotic crossovers. The diploids in our study are homozygous for the *ade2-1* ochre mutation. Diploids with this mutation form red, pink, or white colonies, depending on whether they contain zero, one, or two copies of the *SUP4-o* ochre suppressor tRNA gene, respectively. In our experiments, the diploids were heterozygous for the *SUP4-o* that was located near the end of either the left arm of chromosome V or the right arm of chromosome IV. These diploids form pink colonies. However, a crossover between *SUP4-o* and the centromere results in a red/white sectored colony. Note that only half of the crossovers are detected by the sectoring assay. If chromatids 1 and 4, and 2 and 3, cosegregate, a sectored colony is not formed.

water and then diluted the suspension to a concentration of ~500–1500 cells per plate. After 3 days of growth on MAB6 solid medium at room temperature, the colonies were screened for red/white sectors. We used standard protocols for transformation and DNA isolation.

#### SNP microarray analysis

LOH events were analyzed by SNP microarrays as described previously (St. Charles *et al.* 2012; St. Charles and Petes 2013). These arrays (produced by Agilent) contain four homolog-specific SNPs (W303a or YJM789 specific and Watson or Crick specific) for each of 13,000 SNPs distributed throughout the genome. About 1% of the genome (primarily repetitive regions near the telomere) is not represented on the microarray. The coordinates and sequences of the oligonucleotides on the arrays are described in St. Charles *et al.* (2012) and St. Charles and Petes (2013). DNA was isolated from each sector as described by St. Charles *et al.* (2012). The details for labeling DNA samples, hybridizing them to the SNP microarray, and analyzing the resulting hybridization patterns are provided in detail in St. Charles *et al.* (2012). In brief, DNA from the experimental strain (for example, a white sector) was labeled with Cy5-dUTP, and DNA from the control strain JSC24-2 heterozygous for all SNPs (Yin and Petes 2013) was labeled with Cy3-dUTP. The two samples were mixed and hybridized to the SNP microarray. The microarrays were scanned at wavelengths of 635 and 532 nm, using the GenePix scanner and the GenePix software. The resulting information was exported to text files and examined by R software. The critical data were the ratio of hybridization for the experimental and control samples for each SNP on the array. A ratio of ~1:1 for the experimental and control samples to the four nucleotides representing a particular SNP indicates heterozygosity. A ratio of ~1.5:0.2 or 0.2:1.5 indicates LOH for the SNP. For example, if there is an LOH event in which W303a-derived sequences were duplicated, both W303a-specific oligonucleotides would have a high ratio of hybridization compared to the control, and both YJM789-derived oligonucleotides would have a low ratio.

#### Statistical analyses

Fisher's exact tests and chi-square tests were done using Excel or the VassarStat Web site (<http://vassarstats.net/>). The Mann–Whitney test was done using the R “wilcox.test” function. We calculated 95% confidence intervals using table B11 of Altman (1991).

## Results

### Deletion of *EXO1* reduces the level of UV-induced crossovers, but elevates the level of spontaneous crossovers

In previous studies, we measured the frequency of UV-induced crossovers in a wild-type diploid with the *SUP4-o* gene on chromosome V [PG311 (Yin and Petes 2013)]; the distance between *CEN5* and *SUP4-o* is ~120 kb. A UV dose



**Table 1 Numbers of different types of unselected LOH events in wild-type and *exo1* strains treated with UV**

Strain (genotype)	CO (corrected)	BIR	Conversions unassociated with CO (corrected)	Chromosome loss	Translocation <sup>a</sup>	Interstitial deletion	Total
PG311 (wild type)	60 (120)	21	300 (240)	0	0	0	381
YYy29.8 ( <i>exo1</i> )	11 (22)	5	50 (39)	1	1	1	69

We show the different types of unselected LOH events in the sectored colonies treated with 15 J/m<sup>2</sup> in G<sub>1</sub>-synchronized cells. We examined 16 sectored colonies of YYy29.8 and 47 sectored colonies of PG311. The proportions of CO (crossover), BIR, and conversions unassociated with CO do not differ significantly in the *exo1* and wild-type strains. As described in the text, only half of reciprocal crossovers are detectable because segregation of two recombined chromosomes into one daughter cell and two unrecombined chromosomes into the other daughter is undetectable. In addition, if there is a gene conversion event associated with the crossover following this type of segregation, it would result in an interstitial LOH event, mimicking a gene conversion. Thus, we correct the number of gene conversion events unassociated with a crossover by subtracting the number expected for "hidden" crossovers.

<sup>a</sup> One of the strains derived from YYy29.8 had an unselected terminal duplication of chromosome I sequences (SGD coordinates 181,850–184,217) from the YJM789-derived homolog and a terminal deletion of chromosome XI sequences (SGD coordinates 576,427–578,510) from the W303a-derived homolog. In our related studies (Argueso *et al.* 2008; McCulley and Petes 2010), strains with this LOH pattern had nonreciprocal translocations, reflecting a break in one chromosome with loss of the centromere-distal fragment and repair of the centromere-containing portion of the broken chromosome by a BIR event involving a nonhomolog (generating the duplication). Since we did not confirm the existence of a translocation by other methods, our assignment of this event as a translocation is tentative.

of 15 J/m<sup>2</sup> resulted in 68 red/white sectored colonies from a total of 7194 colonies ( $9.5 \pm 2.2 \times 10^{-3}$ ,  $\pm$  indicating 95% confidence limits). As discussed above, this frequency represents about half the rate of *CEN5-SUP4-o* crossovers and is ~8500-fold higher than the rate observed in untreated cells. The isogenic *exo1* diploid YYy29.8 had a sectoring frequency of  $5.5 \pm 1.9 \times 10^{-3}$  (32 sectored colonies of 5840 total). This reduction is significant by Fisher's exact test ( $P = 0.01$ ). We also measured the frequency of UV-induced sectored colonies in isogenic wild-type (JSC25) and *exo1* (YYy34) strains in which the *SUP4-o* marker was near the telomere of chromosome IV. The frequencies of crossovers in this 1.1-Mb interval were  $9.0 \pm 2.4 \times 10^{-2}$  [127 sectored colonies of 1420 total (Yin and Petes 2013)] and  $3.6 \pm 1.0 \times 10^{-2}$  (52 sectored colonies of 1465 total) in the wild-type and *exo1* strains, respectively. This reduction is significant with a  $P$ -value of 0.0001.

Although the absence of *Exo1p* reduced the frequency of UV-induced crossovers, the rate of spontaneous crossovers was elevated by the *exo1* mutation about threefold. On the right arm of chromosome IV, the sectoring frequencies were  $3.1 \pm 1.0 \times 10^{-5}$  (55 sectored colonies of 1,761,664) and  $1.3 \pm 0.5 \times 10^{-4}$  (23 sectored colonies of 178,055) for the wild-type (St. Charles and Petes 2013) and *exo1* mutant strains, respectively. This elevation in frequency by the *exo1* mutation is significant ( $P = 0.0001$ ).

#### Analysis of classes of LOH events in UV-treated *exo1* diploids

To distinguish among the various classes of LOH events (Figure 1) induced by UV (15 J/m<sup>2</sup>) in the *exo1* diploid

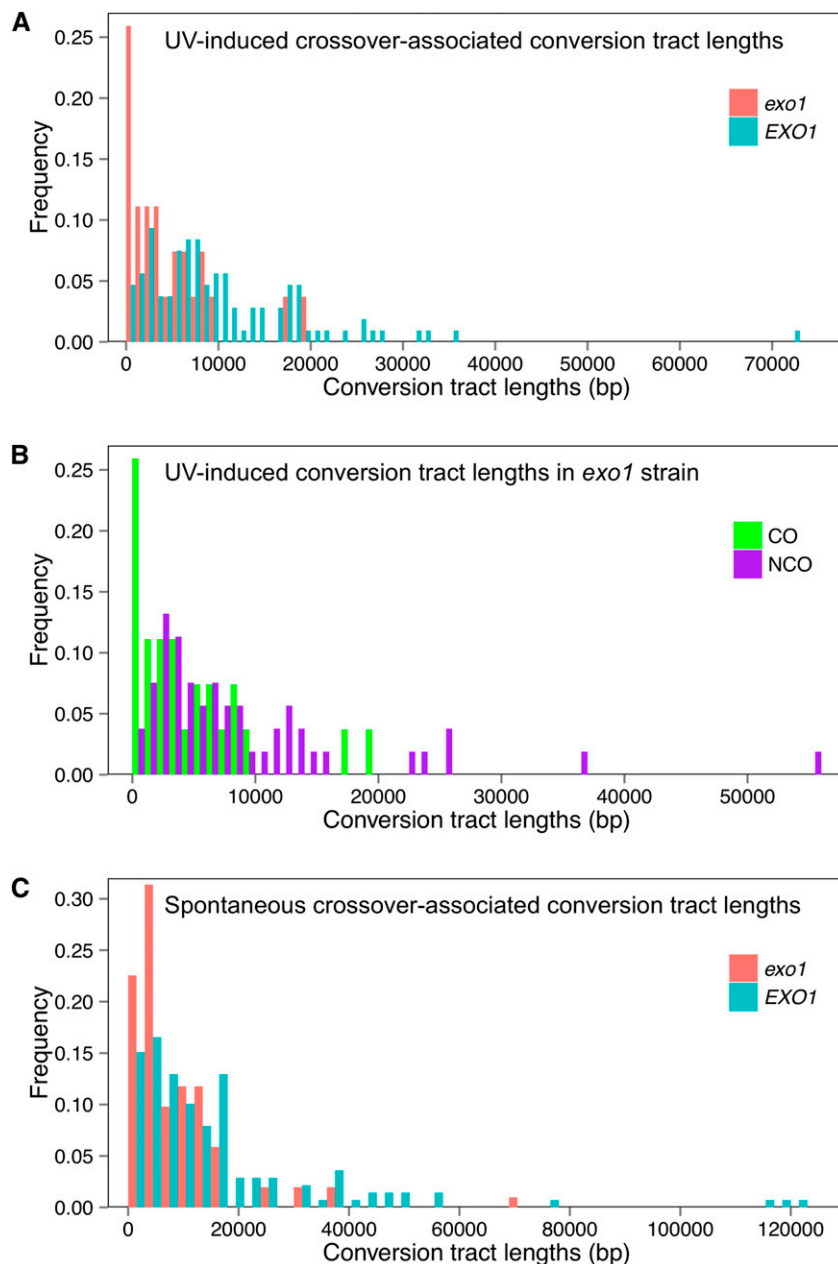
YYy29.8, we mapped LOH events in both sectors of 16 red/white sectored colonies. We used microarrays that assay LOH for the entire genome, allowing us to diagnose the selected event on the left arm of chromosome V as well as unselected events on other chromosomes. Although screening for red/white colonies excludes the possibility of observing BIR on chromosome V, we can detect BIR for unselected events. In addition to the selected event on V, we observed an average of 4.3 unselected LOH events per sectored colony. This frequency is reduced from that observed in a wild-type diploid irradiated with the same UV dose [8.1 LOH events per colony (Yin and Petes 2013)], as expected from the reduced frequency of selected crossovers described above. In Table S1, we depict all LOH events (selected and unselected) derived from UV treatment of G<sub>1</sub>-synchronized YYy29.8 cells. Table S2 has the *Saccharomyces* Genome Database (SGD) coordinates of the transitions between heterozygous and homozygous SNPs for these events.

The numbers of unselected LOH events in the crossover (terminal reciprocal LOH), BIR (terminal nonreciprocal LOH), gene conversion unassociated with a crossover (interstitial LOH), chromosome loss (LOH for one homolog), translocation (terminal duplication or deletion), and interstitial deletion categories are in Table 1 for the wild-type and *exo1* strains; data for the wild-type strain are from Yin and Petes (2013). The distribution of crossovers, BIR, and gene conversion events in the two strains are not significantly different ( $P = 0.77$ ). In Table 1, we also show the numbers of LOH events after correcting for undetectable outcomes (Table 1 legend). If we separate the LOH events into those that involve recombination between homologs and those that do not

**Table 2 Numbers of simple crossovers and SCB and DSCB events in wild-type and *exo1* strains treated with UV**

Strain (genotype)	Simple crossovers	SCB with CO	DSCB with CO	SCB without CO	DSCB without CO	Total
PG311 (wild type)	12	35	57	167	133	404
YYy29.8 ( <i>exo1</i> )	12	10	5	31	19	77

We summarize selected and unselected events detected by the analysis of the genomes of sectored colonies induced by UV. There were a total of 27 crossovers in YYy29.8, 16 selected crossovers on chromosome V and 11 unselected. Of the 104 crossovers in PG311, 44 were selected on chromosome V and the remainder were unselected. The data for PG311 are from Yin and Petes (2013). The determination of SCB and DSCB events was based on the patterns of gene conversion (Table S1). In general, 3:1 conversions are classified as SCBs and 4:0 or 3:1/4:0 hybrid tracts are classified as DSCBs.



**Figure 3** Conversion tract lengths in wild-type and *exo1* strains. Based on the location of LOH regions, we measured conversion tract length in *exo1* strains, both spontaneous events and events induced by UV. These data were compared with measurements performed in wild-type strains (St. Charles and Petes 2013; Yin and Petes 2013). (A) Comparison of UV-induced crossover-associated gene conversion tracts in wild-type and *exo1* strains. A total of 107 and 27 conversion tracts were examined in the wild-type and *exo1* strains, respectively. (B) Comparison of UV-induced conversion tracts in the *exo1* strain. Twenty-seven of the conversion tracts were crossover associated and 54 were unassociated with crossovers. (C) Comparison of spontaneous crossover-associated conversion tracts in wild-type and *exo1* strains. One hundred two conversion tracts were analyzed in the *exo1* strain and 139 were examined in the wild-type strain.

(chromosome loss, translocations, and interstitial deletions), there are significantly more of the second class in the *exo1* strain than in the wild-type strain ( $P = 0.003$ ).

As described in the Introduction, the patterns of LOH observed in sectorized colonies can usually be classified as events involving one broken sister chromatid (SCB) or two broken sister chromatids (DSCB). One event that cannot be assigned as an SCB or a DSCB is a crossover unassociated with gene conversion. We refer to such events as “simple crossovers.” Based on the patterns shown in Table S1, we show in Table 2 the numbers of simple crossovers and SCB and DSCB events, following UV treatment of the *exo1* strain YYy29.8 and the wild-type strain PG311 (Yin and Petes 2013); Table 2 includes both selected and unselected events. The ratio of SCB to DSCB events was not signifi-

cantly different in the wild-type and *exo1* strains ( $P = 0.07$ ). The number of simple crossovers relative to the other classes, however, was significantly elevated in the *exo1* strain relative to wild type ( $P = 0.0003$ ). This finding is likely related to the shorter conversion events associated with crossovers in the *exo1* strain as discussed below.

The data of Table S2 can be used to calculate gene conversion tract lengths for the UV-treated strains. For interstitial LOH events, the tract length was calculated as the average of the maximum tract length (the distance between the heterozygous SNPs flanking the LOH region) and the minimum tract length (the distance between the first and last homozygous SNP in the conversion tract). For conversions associated with crossovers, the length was the average between the maximum tract length (the distance between the last

heterozygous SNP before the conversion tract and the SNP that has the reciprocal LOH pattern indicative of the crossover) and the minimum length (the distance between the first and last homozygous SNPs of the tract). For crossovers with no detectable conversion, we estimated the length of the conversion tract as half the distance between the heterozygous and homozygous SNPs most closely flanking the crossover transition. The median lengths of the crossover-associated conversions for the wild-type and *exo1* strains were 7.6 kb [95% confidence limits of 6.4–9.6 kb (Yin and Petes 2013)] and 3.2 kb (1.3–6 kb), respectively; the *P*-value for the comparison of these lengths is  $8 \times 10^{-5}$  (Mann–Whitney test). Figure 3A is a histogram of these data.

In previous studies of UV-induced recombination in a wild-type strain, we found that the conversion tracts associated with crossovers were significantly longer than those that were unassociated with crossovers, 7.6 kb (6.4–9.6 kb) and 4.8 kb (4.1–5.6 kb), respectively [ $P = 1 \times 10^{-6}$  (Yin and Petes 2013)]. Unexpectedly, in the UV-treated *exo1* strain, the conversion tracts associated with crossovers were significantly ( $P = 0.007$ ) shorter than those unassociated with crossovers, 3.2 kb (1.3–6 kb) and 6.2 kb (3.9–8.5 kb), respectively. Possible explanations for this difference are outlined in the Discussion. Figure 3B is a histogram that shows the distribution of crossover-associated and crossover-unassociated conversion tract lengths in UV-treated *exo1* cells.

#### Analysis of classes of spontaneous LOH events in *exo1* diploids

The frequency of spontaneous sectorized colonies in the *CEN5-SUP4-o* interval in wild-type strains is very low,  $\sim 1 \times 10^{-6}$  (Lee *et al.* 2009). Consequently, we examined spontaneous LOH events in the *exo1* strain YYy34 that has the *SUP4-o* marker near the end of chromosome IV, allowing detection of events in a 10-fold larger interval. Since unselected spontaneous events are rare (St. Charles *et al.* 2012), we restricted our analysis to the selected event on IV by using chromosome IV-specific SNP microarrays (St. Charles and Petes 2013). We examined recombination in 102 red/white sectorized colonies. The depictions of these LOH events are in Table S3 with the coordinates specifying recombination breakpoints in Table S4.

The numbers of simple crossovers and SCB and DSCB events in YYy34 and the isogenic wild-type strain JSC25 are in Table 3. As observed for the UV-induced events, the ratio of SCBs and DSCBs is not significantly different for YYy34 and JSC25 ( $P = 0.06$ ), but the ratio of simple crossovers to other events is significantly higher in the *exo1* strain ( $P = 0.01$ ). This difference is consistent with the observation that the crossover-associated conversion tracts in the *exo1* strain were about half the size of the conversion tracts in the wild-type strains, 5.5 kb (4.5–7.3 kb) and 10.6 kb [8.2–13.6 kb (St. Charles and Petes 2013)], respectively ( $P = 3 \times 10^{-5}$ ). Short conversion tracts would be less likely to be detected using the SNP microarrays and, therefore, would be expected to result in more simple crossovers. The comparison of the lengths of conversion tracts associated with spon-

taneous crossovers for the wild-type and *exo1* strains is in Figure 3C.

Figure 4 shows the location of spontaneous recombination events on chromosome IV in the wild-type (Figure 4A, data from St. Charles and Petes 2013) and *exo1* (Figure 4B) strains. In Figure 4, the y-axis indicates the number of times SNPs on the right arm of chromosome IV were involved in a recombination event, and the x-axis shows the SGD coordinates of SNPs. The peak marked HS4 (hotspot 4) in the wild-type strain is an inverted pair of 6-kb retrotransposons (Ty elements) located between SGD coordinates 981 and 993 kb that is involved in  $\sim 7\%$  of the crossovers in wild type. Recombination at HS4 is significantly ( $P = 0.02$ ) reduced in the *exo1* strain. In addition, there is less recombination at HS3 (a second pair of closely spaced retrotransposons located between SGD coordinates 872 and 884 kb) in the *exo1* strain compared to wild type, although this reduction is not statistically significant. Below, we present models that are relevant to the role of *Exo1p* in generating recombinogenic DNA lesions at HS3 and HS4.

## Discussion

Our studies show several interesting phenotypes in strains that lack *Exo1p*, including (1) a reduction of UV-induced crossovers but an elevation in spontaneous crossovers, (2) loss of a spontaneous crossover hotspot caused by inverted retrotransposons, and (3) conversion events associated with crossovers are shorter than those unassociated with crossovers. Each of these properties is discussed below. All of the UV-induced recombination events examined in our study were derived from cells selected to have a crossover on chromosome V. Although we cannot exclude the possibility that the selection of such sectorized colonies influences the types of events that we recover in UV-treated cells, the types of spontaneous recombination events analyzed in wild-type strains selected for crossovers on two different homologs (chromosomes V and IV) were very similar (Lee *et al.* 2009; St. Charles and Petes 2013).

#### Effect of the *exo1* mutation on the rate of UV-induced and spontaneous crossovers

Although loss of *Exo1p* reduced the frequency of UV-induced crossovers two- to threefold in our assay, the relative frequencies of crossovers, BIR events, and conversions were similar to those observed for wild type (Table 1). In addition, as observed previously for wild-type cells (Yin and Petes 2013), a substantial fraction (about one-third) of the UV-induced conversion events in *exo1* strains reflected repair of two sister chromatids broken at approximately the same position (DSCB events). We interpret this result as indicating that high doses of UV in G<sub>1</sub>-synchronized cells result in recombinogenic DSBs (Yin and Petes 2013). We emphasize that other types of UV-related DNA lesions may be more common than DSBs in unreplicated chromosomes. For example, DNA molecules with single-stranded nicks could be replicated to generate single chromatid breaks



**Table 3** Numbers of spontaneous simple crossovers and SCB and DSCB events in wild-type and *exo1* strains

Strain (genotype)	Simple crossovers	SCB with CO	DSCB with CO	Total
JSC25 (wild type)	18	31	90	139
YYy34 ( <i>exo1</i> )	27	29	46	102

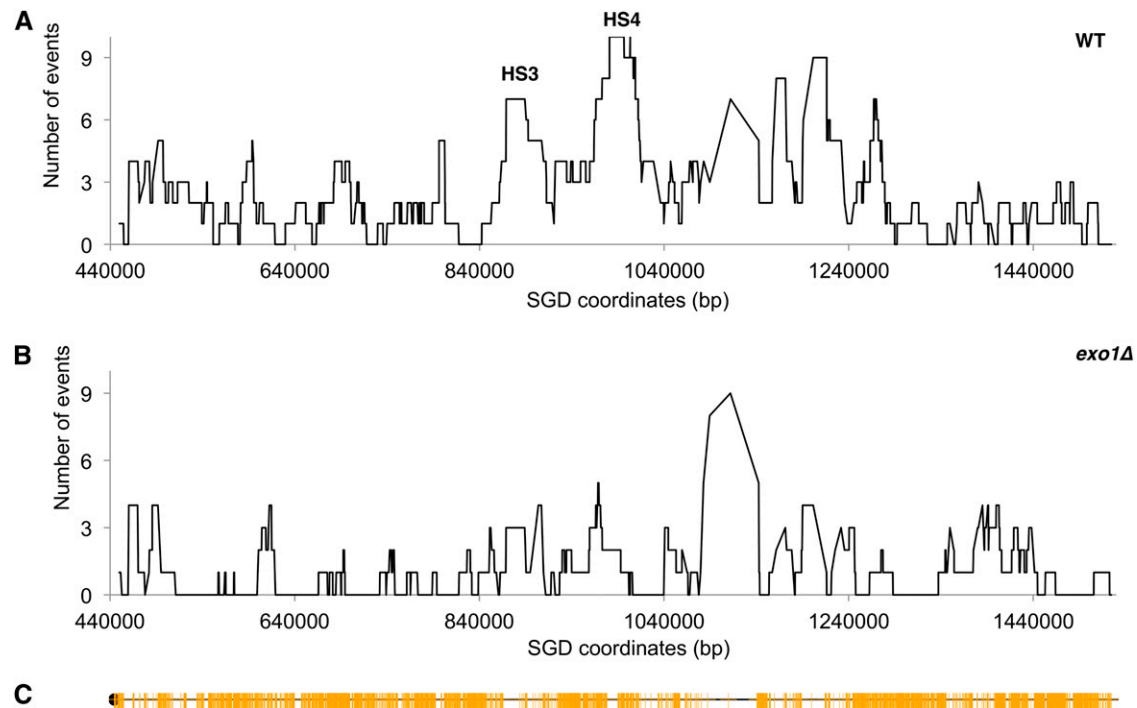
We summarize the analysis of conversions associated with spontaneous crossovers in JSC25 and YYy34. The data for the wild-type strain are from St. Charles and Petes (2013). The determination of SCB and DSCB events was based on the patterns of gene conversion in Table S3.

(Galli and Schiestl 1999). Such lesions are likely to be responsible for the SCB events that we observed. Based on previous studies (Kadyk and Hartwell 1992), many of the SCB lesions are usually repaired by sister-chromatid recombination, an event that is not detectable by our analysis.

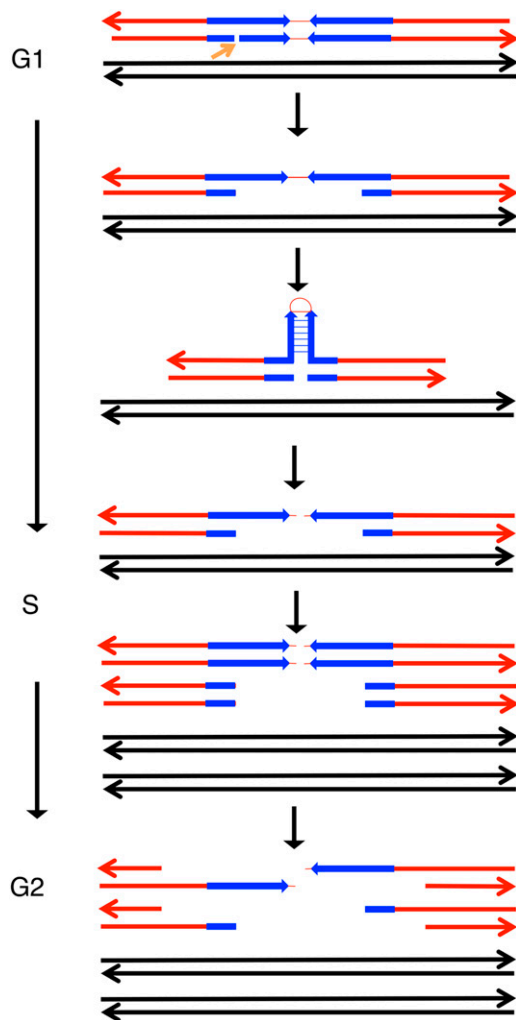
On the basis of previous results (Giannattasio *et al.* 2010; Ma *et al.* 2013; Yin and Petes 2013), we proposed several mechanisms by which *Exo1p* would have a role in production of UV-associated DSBs in G<sub>1</sub>-synchronized cells (Yin and Petes 2013). One possibility is that the DSB reflects two closely spaced 30-base gaps resulting from removal of dimers on opposite strands. Expansion of one of these short gaps into a longer gap that overlaps with the gap on the opposite strand would result in a DSB. Previously, we calculated that a dose of 15 J/m<sup>2</sup> would result in ~35 closely opposed ( $\leq 75$  bp) pairs of lesions, assuming a random distribution (Yin and Petes 2013). However, Lam and Reynolds (1987) observed that UV-induced lesions <15 bases apart were observed at a frequency that was

higher than expected for a random distribution. By either estimation, there appear to be a sufficient number of closely opposed lesions to generate the observed recombination events.

The reduction in the frequency of UV-induced crossovers in our system in the *exo1* mutant could reflect either a reduction in the frequency of recombinogenic lesions or less efficient processing of these lesions into a crossover between homologs. Since all types of recombination events (crossovers, noncrossovers, and BIR events) are reduced by the *exo1* mutation by approximately the same extent, we favor the possibility that recombinogenic lesions are reduced by loss of *Exo1p*, although we cannot rule out the second possibility. The observation that loss of *Exo1p* reduces, but does not eliminate, mitotic crossovers could reflect redundancy in gap processing by the *Sgs1p/Top3p/Rmi1p* and *Dna2p* pathway. Ma *et al.* (2013) interpreted the reduction in sister-chromatid recombination events observed in UV-treated G<sub>2</sub>-synchronized *exo1 rad30* cells as



**Figure 4** Distribution of spontaneous crossovers on the right arm of chromosome IV in wild-type and *exo1* strains. We show the frequency with which SNPs on the right arm of chromosome IV were involved in a conversion/crossover in the strains JSC25 [wild type (St. Charles and Petes 2013)] and YYy34 (*exo1*). More than 100 crossovers were mapped in each strain. (A) Crossovers and associated conversions in the wild-type strain. HS3 and HS4 are hotspots for recombination, each containing inverted pairs of Ty elements. (B) Crossovers and associated conversions in the *exo1* strain. (C) Location of SNPs used in mapping. Each yellow line shows a SNP position. Note that there is a region of ~65 kb that contains only one SNP.



**Figure 5** Role of Exo1p in regulating HS4 hotspot activity. The HS4 G<sub>1</sub>-specific hotspot for spontaneous mitotic recombination contains two closely linked Ty elements. Since deletion of one of the elements or expansion of the distance between the elements results in loss of hotspot activity, we previously suggested that the recombinogenic effects of HS4 likely involve the formation of a hairpin or cruciform structure (St. Charles and Petes 2013). The two Ty elements of HS4 (shown in blue) are present on the W303a-derived homolog (red) but not the YJM789-derived homolog (black). One scenario for production of a DSB in G<sub>1</sub> is that a nick on one strand of the inverted repeat (shown by the arrow) is expanded into a gap by Exo1p, allowing formation of a hairpin. A subsequent nick of the sequences at the tip of the hairpin would produce a DSB. The resulting broken ends would require extensive resection (another function of Exo1p) to allow pairing with the YJM789-derived homolog.

a reduction in recombinogenic lesions; in their experiments, however, they suggested that the recombinogenic lesion was a large single-stranded gap rather than a DSB. In meiosis, Exo1p affects both resection of Spo11p-generated chromosome ends and resolution of double Holliday junctions into crossovers without an effect on noncrossovers (Zakharyevich *et al.* 2010). Since we see a reduction in crossovers, noncrossovers, and BIR events, the reduction in UV-induced mitotic events is not likely to reflect the same resolution role as observed in meiosis.

In contrast to the reduction in UV-induced crossovers, we observed an elevation in the frequency of spontaneous crossovers in the *exo1* strain relative to the wild type. This result argues that Exo1p is not required for processing spontaneous DNA damage into recombinogenic lesions. Because of the multiple cellular roles of Exo1p and the possibly diverse sources of spontaneous DNA damage, we cannot assign a unique explanation for the modest hyper-Rec phenotype. One possibility is that the reduction reflects the role of Exo1p in inhibiting recombination between diverged DNA sequences (Nicholson *et al.* 2000). Alternatively, since Exo1p mediates an error-free pathway in postreplication repair (Karras *et al.* 2013), loss of Exo1p could channel more lesions into inter-homolog exchanges.

#### Genomic locations of UV-induced and spontaneous recombination events in *exo1* strains

The map positions (SGD coordinates) of UV-induced events in the *exo1* strain Y<sub>Y</sub>29.8 are given in Table S2. The events are distributed widely over the chromosomes with no strong hotspots, similar to observations of UV-induced recombination events in wild-type strains (Yin and Petes 2013). All chromosomes have at least one event, and the number of events per chromosome is correlated with the size of the chromosome ( $r^2 = 0.73$ ;  $P = 0.001$ ). We also analyzed the LOH regions to find out whether various chromosome elements [for example, centromeres, transfer RNA (tRNA) genes, G-quadruplex sequences, etc.] were overrepresented at the breakpoints between the heterozygous markers immediately flanking the LOH regions for conversions unassociated with crossovers and between the heterozygous and homozygous regions for crossovers; these breakpoints are likely to include the regions of the chromosome that underwent the recombinogenic DNA lesion. The details of the analysis are summarized in File S1, and the results are in Table S5. After correction of the *P*-values for multiple comparisons (Hochberg and Benjamini 1990), none were significant at a value of 0.05.

We also examined the distribution of spontaneous recombination events in the *exo1* strain Y<sub>Y</sub>34 on the right arm of chromosome IV. The most obvious difference between the wild-type and *exo1* strains in this distribution was the absence of the HS3 and HS4 peaks in the *exo1* strain (Figure 4). The HS3 and HS4 hotspots in the wild-type strain have two important properties: they are G<sub>1</sub> specific and they involve breakage of the W303a-derived homology (St. Charles and Petes 2013). We found that the W303a-derived homolog has an inverted pair of Ty elements spaced ~50 bp apart at HS4, whereas the YJM789-derived homolog has only a portion of one Ty element at the allelic position (St. Charles and Petes 2013). Since deletion of one of the Ty elements or increasing the distance between the elements results in loss of HS4 activity in the wild-type strain, we suggested that the recombinogenic effect of HS4 likely reflected formation of a secondary structure in unreplicated DNA (St. Charles and Petes 2013). One mechanism for

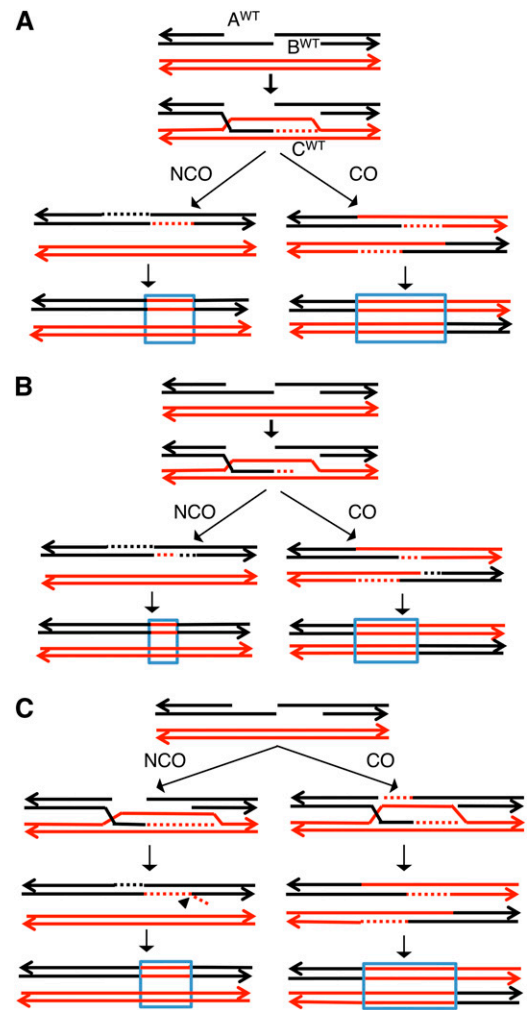
producing a DSB in  $G_1$  is to expand a nick on one strand into a gap, allowing for formation of a “hairpin” on the ungapped strand (Figure 5). Cleavage of the unpaired bases at the tip of the hairpin would result in a  $G_1$ -specific DSB. The role of *Exo1p* in this process could be in the formation of the gap and/or the processing of the broken ends to allow invasion of the homolog that lacks Ty elements. An interesting feature of the conversion events associated with HS4 in wild-type strains is that median conversion tract length is  $\sim 50$  kb, much larger than that observed for other spontaneous  $G_1$  gene conversions [14 kb (St. Charles and Petes 2013)].

We also examined the spontaneous crossovers in the *exo1* strain for over- or underrepresentation of chromosome elements (Table S5). Although no element was significantly elevated, we found a significant underrepresentation of tandem repeats in the LOH regions in the *exo1* strain. A similar underrepresentation was found previously for spontaneous crossovers in the wild-type strain (St. Charles and Petes 2013). Chromosome IV has 98 different tandem repeats (Tandem-repeat Database; <https://tandem.bu.edu/cgi-bin/trdb/trdb.exe?taskid=0>) with repeat units varying in size between 1 and 213 bp with the number of repeats per array varying between 2 and 33 copies. It is possible that a subset of these repeats has a chromatin modification or other property that renders them resistant to recombinogenic DNA lesions.

#### Relationship between resection of DNA ends and the length of conversion tracts

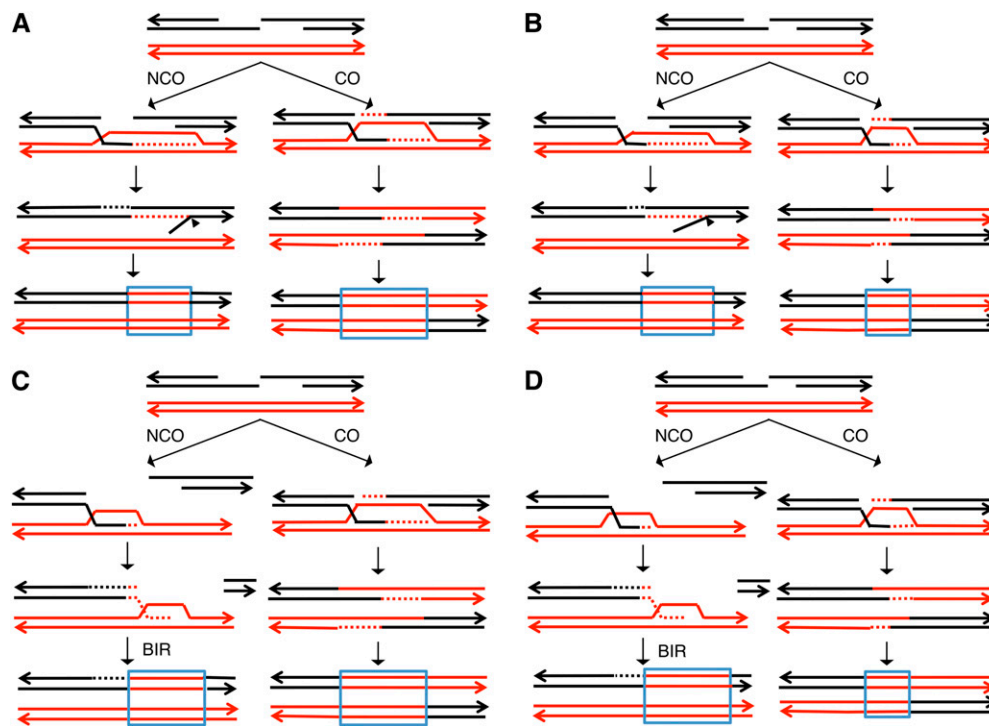
In depictions of the double-strand break repair model, the extent of the gene conversion tract is usually shown as equivalent to the extent of end resection (Figure 6A). For a recombination event in a wild-type strand, we indicate the resection length of the invading chromosome end as  $A^{WT}$  and the resection length of the other end as  $B^{WT}$ , and the simplest assumption is that  $A^{WT} = B^{WT}$ . Further, we hypothesize that amount of DNA synthesized by the invading end ( $C^{WT}$ ) is equivalent to the resection length. With these assumptions, one would expect that the length of gene conversion tracts unassociated with crossovers would be half of that associated with crossovers. In addition, we would expect both types of conversion tracts would be shorter in the *exo1* strain because of less resection. If  $A^{WT}$  and  $B^{WT}$  are not equal, nonetheless, the conversion tracts unassociated with crossovers would be expected to be shorter than those associated with crossovers.

A number of studies have shown that the relationship between end resection and conversion tract length is more complicated than illustrated in Figure 6A. For example, although *mre11* strains (which have reduced extents of end resection) have reduced conversion tract lengths in plasmid–chromosome gap-repair assays (Symington *et al.* 2000; Krishna *et al.* 2007), conversion tract lengths involving homologs are unaffected by the *mre11* mutation (Krishna *et al.* 2007). In addition, the *yku70* mutation, which leads to increased resection, does not result in longer



**Figure 6** Patterns of crossover-associated and crossover-unassociated conversion tracts based on current models of recombination. Black and red lines show DNA strands of the two homologs with arrows marking the 3' ends. Dotted lines show repair-associated DNA synthesis. For all of the models, we assume that conversion events unassociated with crossovers (NCO) occur as a consequence of SDSA, and events associated with crossovers (CO) reflect resolution of Holliday junctions. We assume that the two broken ends are resected to the same extent ( $A^{WT} = B^{WT}$ ), although our conclusions do not require this assumption (details in text).  $C^{WT}$  shows the length of DNA synthesized by the invading strand. In heteroduplexes with one black strand and one red strand, we show correction of the mismatches to generate two red strands. The resulting conversion tracts are outlined in blue. (A) The length of DNA synthesized by the invading strand ( $C^{WT}$ ) equals the amount of resection, and  $A^{WT} = B^{WT}$ . The expected conversion tract length for events associated with crossovers is about twice that for events unassociated with crossovers. (B) If  $C^{WT} < A^{WT}$  and  $B^{WT}$ , then the relative length of the crossover-associated tract compared to the crossover-unassociated tract is greater than in A. (C) If  $C^{WT} > A^{WT}$  and  $B^{WT}$ , one possibility is that, during the SDSA event, the 3' end of the reinvading strand is displaced. Removal of this end by a branch-processing enzyme such as Rad1p/Rad10p (Mazón *et al.* 2012) would result in a conversion tract in the NCO pathway that is the same length as in A.

conversion tract lengths in recombination events between homologs (Krishna *et al.* 2007). Note that the effect of these mutants on resection could be lesion specific (Symington and Gautier 2011). Finally, in studies of plasmid–chromosome



**Figure 7** Two models for the formation of long gene conversion tracts that are unassociated with crossovers. The data that require explanation are (1) crossover-associated conversion events are shorter in the *exo1* strain than in wild type and (2) crossover-unassociated conversions are longer than crossover-associated conversions in the *exo1* strain. The labels are the same as in Figure 6. The observations can be explained by an oversynthesis model (A and B) or by a BIR model (C and D). (A) UV-induced conversion in the wild-type strain (oversynthesis model). The synthesis from the invading strand is more extensive than the amount of DNA resected in the NCO pathway. The reinvasion during SDSA displaces the 5' end of invaded homolog, and the resulting single-stranded branch is removed (shown by a triangle). In the CO pathway, the heteroduplex region is limited by resection. (B) UV-induced conversion in the *exo1* strain (oversynthesis model). The conversion tract in the NCO pathway is similar to that observed in wild type, whereas the conversion tract for the CO pathway is shorter due to less resection. (C) UV-induced conversion in the wild-type strain (BIR model). In the NCO pathway, the conversion is a consequence of BIR. Following copying of the red chromosome, the invasion is reversed, and the end generated by BIR reassociates with the broken black chromosome. In this model, the conversion tract is not a consequence of repair of mismatches in a heteroduplex. The conversion events in the CO pathway occur by the same mechanism as in A and B. (D) UV-induced conversion in the *exo1* strain (BIR model). As in C, the BIR event generating the conversion tract in the NCO pathway is not limited by resection, unlike the conversion tract in the CO pathway.

served in wild type, whereas the conversion tract for the CO pathway is shorter due to less resection. (C) UV-induced conversion in the wild-type strain (BIR model). In the NCO pathway, the conversion is a consequence of BIR. Following copying of the red chromosome, the invasion is reversed, and the end generated by BIR reassociates with the broken black chromosome. In this model, the conversion tract is not a consequence of repair of mismatches in a heteroduplex. The conversion events in the CO pathway occur by the same mechanism as in A and B. (D) UV-induced conversion in the *exo1* strain (BIR model). As in C, the BIR event generating the conversion tract in the NCO pathway is not limited by resection, unlike the conversion tract in the CO pathway.

recombination in an assay involving 800 bp of homology, Guo and Jinks-Robertson (2013) found that conversion tracts unassociated with crossovers had the same lengths in wild-type and *exo1 sgs1* strains.

Variants of the model shown in Figure 6A in which the length of DNA synthesized by the invading end is not identical to the length of resection have been recently described (Mazón *et al.* 2012; Guo and Jinks-Robertson 2013). In Figure 6B, we show a model in which the invading strand disassociates from the template chromosome after synthesizing a segment that is shorter than the gap resulting from resection. By this model, the conversion tracts unassociated with crossovers would be shorter than those associated with crossovers as in Figure 6B, but the lengths of the conversion tracts would not necessarily be affected by the *exo1* mutation. This model, however, does not explain our analysis of UV-induced events. In the wild-type strain, the conversion tracts unassociated with crossovers (4.8 kb) are greater than half the size of those associated with crossovers (7.6 kb), and the conversion tracts unassociated with crossovers in the *exo1* strain (6.2 kb) are longer than those associated with crossovers (3.2 kb).

It was shown recently that the amount of DNA synthesized from the invading end could exceed the length of the resected gap (Mazón *et al.* 2012). The subsequent Synthesis-Dependent Strand Annealing (SDSA) event leads to a single-stranded 3' "tail" that was processed by the Rad1–Rad10

nuclease (Figure 6C). By this mechanism, the extent of the heteroduplex would be regulated by the length of resection as in Figure 6A. Similar to the other models in Figure 6, this model is not consistent with our observations in the *exo1* strain. One difference between the experiments of Mazón *et al.* and our experiments is that the system used by Mazón *et al.* involves the interactions of ectopic repeats in which flanking heterology limits the size of the conversion tract.

We propose two new models to explain our data (Figure 7). Both require the assumption that the conversion events associated with crossovers are fundamentally different from those unassociated with crossovers. We suggest that the conversion events associated with crossovers are limited by the length of the resection tract, consistent with the observation that the conversion events associated with crossovers in wild type are significantly longer than those observed in the *exo1* strain. In contrast, for the models shown in Figure 7, the lengths of the conversion tracts that are unassociated with crossovers are not limited by resection. In the first model (Figure 7A, wild type; and Figure 7B, *exo1*), the invading strand synthesizes a region of DNA longer than the gap resulting from end processing. When the invading strand is displaced by SDSA, the displaced end reinvasades the processed black chromosome, producing a displaced 5' end. This end could be removed by Rad27p or a related flap-processing enzyme. By this model, the conversion



events unassociated with crossovers can be longer than conversions associated with crossovers (Figure 7B). The processing of a 5' flap was previously proposed as a step in meiotic recombination (Osman *et al.* 2003), although in this model, the flap was generated during resolution of a recombination intermediate into a crossover rather than as a step in producing a conversion event unassociated with a crossover.

For the alternative model (Figure 7C, wild type; and Figure 7D, *exo1*), crossover-associated conversions follow the same pathway as in Figure 7, A and B. The conversions that are unassociated with crossovers, however, are generated by BIR. Following conservative DNA synthesis (Donnianni and Symington 2013; Saini *et al.* 2013) of a segment of the invaded chromosome, we suggest that the invading DNA molecule dissociates, rejoining the broken end. To be consistent with our data, the length of the conversion tract formed by BIR must be longer than the length of the processed ends. The primary difference between conversion tracts of the two models in Figure 7 is that the tracts shown in Figure 7, A and B, are a consequence of mismatch repair in long heteroduplexes whereas those of Figure 7, C and D, result from conservative DNA synthesis during BIR (Donnianni and Symington 2013; Saini *et al.* 2013).

Although the suggestion that the heteroduplex regions of crossover-associated and crossover-unassociated conversion events are formed differently may appear problematical, studies of meiotic recombination events in yeast provide a precedent. Meiotic conversion events of the two types are under different genetic regulation (Allers and Lichten 2001; Börner *et al.* 2004) and involve the resolution of different types of intermediates. In summary, our observations of UV-induced gene conversion events in wild type and *exo1* strongly suggest a fundamental difference in mechanisms by which conversion tracts are generated for crossover and noncrossover pathways.

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# GENETICS

Supporting Information

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## **The Role of Exo1p Exonuclease in DNA End Resection to Generate Gene Conversion Tracts in *Saccharomyces cerevisiae***

Yi Yin and Thomas D. Petes

## Associations of LOH breakpoints with various elements of chromosome structure

In chromosomes with LOH events, the regions of transition likely contain the sites of the recombinogenic DNA lesion. Consequently, for each recombination event listed in Tables S2 and S4, we assigned “windows” likely to contain the initiating lesion; the method that we used is described in detail in Yin and Petes (2013). In brief, the window was defined from the leftmost coordinate of the first transition to the rightmost coordinate of the last transition. For example, in sector 1R/W, for the hybrid conversion tract on chromosome VII, the window is from SGD coordinates 663893 to 686220.

The *exo1* data for UV-induced recombination events (Table S2) was analyzed separately from the spontaneous events (Table S4). For each of the two datasets, we summed the sizes of all of the recombination windows. For the UV-induced samples, we restricted the analysis to unselected recombination events (those that did not occur on the left arm of chromosome V). This sum was 817,055 bp for these events. We examined 16 sectorized colonies induced by UV, and the amount of genomic DNA (other than the left arm of chromosome V) that is represented on the SNP microarray is about 11.4 Mb. Thus, for the UV-induced events, the fraction of genomic DNA represented in the windows is  $817 \text{ kb} / (11.4 \times 16 \text{ Mb})$  or  $4.48 \times 10^{-3}$ . Based on the number and location of elements in the relevant region of the genome, we can then calculate the expected numbers of elements if they are randomly distributed with respect to the recombination windows. For example, from an examination of the location of palindromic sequences greater than 16 bp in length (Lisnic *et al.*, 2005), we calculate that there are 563 palindromes in the genomic region that does not include the right arm of V. Examining sixteen genomes, therefore, we expect, therefore, that there will be 40 palindromes located in the windows and 8968 located outside the windows (Table S5). We observed almost exactly these numbers (38 in the windows and 8970 outside of the windows). By chi-square analysis, there is no significant difference between the expectation and the observations. The comparable calculations were done using 13 other genetic elements (Table S5).

We performed a similar calculation with the spontaneous recombination events. The sum of the recombination windows for the spontaneous events selection on chromosome IV was 1,259,222 bp. The amount of DNA represented on the right arm of chromosome IV that is on the SNP microarray is 1,074,795 bp, and we examined a total of 109,629,090 bp ( $1,074,795 \text{ bp} \times 102$  sectorized colonies). The windows, therefore, represent about  $1.14 \times 10^{-2}$  of the total DNA examined. We compared the observed

and expected chromosome elements within the windows, and outside of the windows as described above for UV-induced events. Since we performed multiple comparisons, we corrected *p* values using the method of Hochberg and Benjamini (1990). After this correction was applied, only one comparison was significant. Tandemly-arrayed repeats were significantly under-represented in the spontaneous crossovers.

The references for the number and location of various genomic elements are: ncRNAs, snoRNAs, snRNAs, retrotransposons, tRNA genes, centromeres, ARS elements, solo LTRs (SGD Website: <http://www.yeastgenome.org/cgi-bin/seqTools>); G4 quadruplex DNA (Capra *et al.*, 2010); strongly- and weakly-transcribed genes (Nagalakshmi *et al.*, 2008); Rrm3p pause sites (Azvolinsky *et al.*, 2008); gamma-H2AX peaks (Szilard *et al.*, 2010); TER sequences (Fachinetti *et al.*, 2010). The strongly- and weakly-transcribed genes were based on examining the transcription of genes within the analyzed area, and picking the top 5% and bottom 5% levels of transcription, respectively.

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**Tables S1-S6**

Available for download as Excel files at <http://www.genetics.org/lookup/suppl/doi:10.1534/genetics.114.164517/-/DC1>

**Table S1** Classes of LOH events in sectored colonies of the *exo1* strain YYy29.8 treated with 15 J/m<sup>2</sup> of UV

**Table S2** SGD coordinates for LOH transitions in the *exo1* strain YYy29.8 induced by a UV dose of 15 J/m<sup>2</sup> in G1-synchronized cells

**Table S3** Classes of spontaneous reciprocal crossovers in red/white sectored colonies of the strain YYy34

**Table S4** SGD coordinates for transitions in *exo1* spontaneous sectors

**Table S5** Analysis of over- or under-represented genome elements within spontaneous and unselected UV-induced conversion tracts in the *exo1* mutant

**Table S6** Strain construction